

## *In vitro* propagation of *citrus aurantiifolia* cv. Sai Sharbati.

Sherkar Sandip Harishchandra

Genuine Biotech, Gut No 121/2, Sambhaginagar, Dist-Osmanabad MH-413 501

\*Corresponding author Email: [sandiphsherkar@gmail.com](mailto:sandiphsherkar@gmail.com)

### Manuscript Details

Available online on <https://www.irjse.in>

ISSN: 2322-0015

Editor: Dr. Arvind Chavhan

### Cite this article as:

Sherkar Sandip Harishchandra. *In vitro* propagation of *citrus aurantiifolia* cv. Sai Sharbati., *Int. Res. Journal of Science & Engineering*, 2020, Special Issue A9: 53-59.

Article published in Special issue of International e-Conference on "Emerging trends and Challenges In life sciences" organized by Department of Botany, Indraraj Arts, Commerce & Science College, Sillod-431112, Dist Aurangabad, Maharashtra, India date, June 18-19, 2020.



Open Access This article is licensed under a Creative Commons Attribution 4.0 International License, which permits use, sharing, adaptation, distribution and reproduction in any medium or format, as long as you give appropriate credit to the original author(s) and the source, provide a link to the Creative Commons license, and indicate if changes were made. The images or other third party material in this article are included in the article's Creative Commons license, unless indicated otherwise in a credit line to the material. If material is not included in the article's Creative Commons license and your intended use is not permitted by statutory regulation or exceeds the permitted use, you will need to obtain permission directly from the copyright holder. To view a copy of this license, visit <http://creativecommons.org/licenses/by/4.0/>

### Abstract

The present study was undertaken to develop efficient protocol for *in vitro* propagation of *Citrus aurantiifolia* cv. Sai Sharbati through nodal shoot segments. Explants produced multiple buds when cultured on murashige and skoog's medium containing 0.5 mg<sup>l</sup><sup>-1</sup> BAP and 1.0 mg<sup>l</sup><sup>-1</sup> GA<sub>3</sub>. BAP was recorded to be better than kinetin in terms of multiplication rate, average shoot height and average number of leaves. Separated shoots were rooted on MS medium containing different concentrations of growth hormones NAA and IBA, from which half strength medium supplemented with 1.0 mg<sup>l</sup><sup>-1</sup> IBA was superior to NAA. *In vitro* plantlets were hardened using different potting mixtures, highest survival (90.23 %) success was achieved after transfer to cocopeat. *In vitro* plantlets were successfully acclimatized in greenhouse.

**Keywords:** Nodal shoot segments, BAP, IBA, MS medium.

### Introduction

Citrus is an important genus of evergreen fruit crops belongs to family *Rutaceae*, originated in Southeast Asia bordered by Northeastern India and the Yunnan province of China [1, 2]. In India, citrus occupies an area of 2, 83,000 Ha with production of 32, 21,000 MT [3]. It is considered as number one fruit of the world due to its high nutritional value, great production potential and preparation of large number of fruit products from them [4]. *Citrus aurantifolia* is small fruited acid lime mainly used in daily consumption and in juice production. It is widely used throughout the globe because of its various properties like antibacterial, anticancer, antidiabetic, antifungal, anti-hypertensive, anti-inflammation and antioxidant [5].

Acid lime is generally propagated through cutting, air-layering and budding on seedling rootstocks [6]. Propagation of plants is an often difficult, expensive and season specific when buds are available. Success ratio of these methods of propagation is not affordable. Air layering of plants never yields the required quantity of planting materials. These conventional techniques are also not free from risk of in-borne pathogens. Citrus trees also declining on a large scale due to diseases like tristeza, viruses, canker and various other factors; for example 823 different insects species attacked citrus spp. [7]. However, tissue culture methods offer an alternative means of vegetative propagation in short time and can overcome some constraints to citrus improvement and cultivation throughout year with quality fruits and resistance to diseases and environmental stresses [8]. There are very few data available about in vitro propagation of cv. Sai Sharbati.

The present study was conducted with an objective to develop an efficient protocol for in vitro propagation of cv. Sai Sharbati. The studies have focused on establishment of cultures, effects of different cytokinin and auxins on shoot multiplication and in vitro rooting and to standardize the culture medium, which is most important factors for the success of tissue culture of cv. Sai Sharbati.

## Methodology

**Plant material:** Healthy and high quality fruit yielding plants of Citrus aurantiifolia cv. Sai Sharbati were selected as a mother plant from Savita agro farm, Osmanabad (MH). Shoot cuttings were procured from the selected mother plants in the month of September, 2016. These explants were collected in glass bottle containing tap water to prevent wilting until they were bring to laboratory. Leaves were removed from shoot cutting and washed carefully under running tap water to remove micro-flora and dirt before surface sterilization.

**Media preparation:** The stock solutions required for Murashige and Skoog [9], vitamins, growth regulators

were prepared well in advance and stored in refrigerator. During medium preparation, each stock solutions after bringing it to room temperature was added one by one in the required quantity to the beaker containing small amount of distilled water. After addition of sucrose (30 g/l), meso inositol (100 mg/l) and growth regulators, final volume was prepared by adding filtered water. The pH of the medium was adjusted to 5.8 with the help of 1N NaOH and/or 1N HCl. Thereafter, the gelling agent agar was added for solidification of the medium. The medium was then boiled and dispensed in culture glass bottles in equal volume for aseptic manipulations and autoclaved at pressure of 15 lbs /inch<sup>2</sup> and at temperature of 121°C for 20 min for sterilization. The medium was stored in the dark at 25±2°C and used after 3-4 days of sterilization.

### Surface sterilization and inoculation of explants:

Nodal segments excised from the selected healthy tree were used as explants for *in vitro* propagation. Shoot cuttings were immersed in an aqueous solution of liquid detergent containing 2–3 drops of Tween-20 for 10 min and then treated with carbendazime for 30 min. The explants were taken to laminar air flow for surface sterilization. These explants were treated with 70% ethanol for 30 to 40 seconds, followed by different concentration of sterilants like mercuric chloride (HgCl<sub>2</sub>) and sodium hypochlorite (NaOCl) (Table-1). After 3-4 times rinsing with sterile autoclaved water, nodal segments of 2.0–3.0 cm were inoculated on MS medium under laminar air flow.

**Culture conditions:** Inoculated cultures were kept under controlled conditions of temperature (26± 2°C), light for 12 hr photoperiod and 60% relative humidity. Per cent bud establishment was recorded during four weeks of culture. Sprouted buds were then subcultured on multiplication and rooting medium fortified with different concentration of growth regulators. Observations were recorded regularly for shoot growth, fungal/ bacterial contamination and data was recorded after 4 weeks of culturing.

**In vitro multiplication:** Healthy shoots were transferred on MS medium supplemented with different

concentrations and combination of growth hormones like BAP, Kin and GA<sub>3</sub> to investigate their effects on enhancement of shoot multiplication. Nodal and shoot tip explants were used for shoot multiplication. Shoots were multiplied by the method of enhanced release of axillary buds [10]. The observations were recorded after 4 weeks of cultures as- rate of multiplication, Quality of shoots, average shoot length and leaves (Table 2). The multiplication medium which showed healthy, long and good quality micro shoots during incubation period was selected for further multiplication on same medium.

**In vitro rooting:** Good quality micro shoots (20–30 mm long) were used for root induction produced from *in vitro* multiplication stage were individually separated and transferred to full strength and half strengths of MS medium supplemented with various concentrations of IBA or NAA (Table 3). The individual shoot bases were cut transversely. Four such shoots were inoculated in each culture bottle. Three replications of each treatment were made. In all rooting experiments 6 g/l agar was used to easily removal of plantlet during hardening. Data was recorded on Days taken for root initiation, percent rooting, Root length and numbers of roots per shoots after 4 weeks of culturing period.

**Hardening and acclimatization of plantlets:** For hardening and acclimatization, *in vitro* rooted plantlets were carefully removed from the culture glass bottles to avoid any damage to delicate root system and kept in a beaker and washed under running tap water for 30 min to remove agar sticking to root surface. Thereafter, the plantlets were kept dipped in solution of 0.2 % carbendazim for 15-20 minutes before transferring to different potting mixture. Root portion was placed inside the mixture gently to avoid any injury. In order to maintain high relative humidity, plantlets were covered with glass jars and watered at 2-3 days intervals. Pots were kept at temperature of 28±2°C in polytonal. When plants showed initial signs of establishment, humidity was decreased by removing the glass jars for few minutes. Thereafter, the plants were hardened by removing the glass jars for increased time intervals to reduce the relative humidity gradually. Observations

were recorded for survival percentage, average number of leaves and average height of plants (Table 4). Eight weeks old primary hardened plants were transferred to polybags containing soil media (red soil and FYM, 3:1) for secondary hardening before dispatch to field transfer.

**Statistical analysis:** The experiments were repeated three times and subjected to completely randomized design [11]. Significance of treatment on various observations was determined using analysis of variance (ANOVA) technique for CRD. If experimental treatments found significant, their relative performance was tested with critical difference (CD<sub>0.05</sub>).

## Results and Discussions

**Shoot multiplication:** Nodal explants of cv. Sai Sharbati resulted in 64.43 per cent uncontaminated cultures after four weeks of incubation when treated with 0.1% HgCl<sub>2</sub> for 5 min. The results obtained are in conformity with Rana and Singh [12], surface sterilization is necessary to make explants free from all contaminants [13]. These cultured nodal segments showed 74.80 per cent proliferation after 3 weeks of culture on MS basal medium (Table-1; Fig-1). Proliferated shoots were then cut into small pieces (2-3 nodal segments) and cultured on MS medium supplemented with different concentration and combination of growth regulators. Different concentration and combination of growth regulator showed mark influence on shoot multiplication, length of shoots and leaf numbers. All medium combinations showed multiplication of shoots (Table-2). The highest multiplication rate of 1:2.8 was observed in MS medium supplemented with 0.5 mg l<sup>-1</sup> BAP and 1.0 mg l<sup>-1</sup> GA<sub>3</sub> (Table-2 and Figure-2). Similarly, the highest shoot length (4.53 cm) and average number of leaves (5.03) was observed on this medium as compare to other. Similar study of Sujata *et al.*, 2010[14] showed, BAP was best cytokinin for citrus shoot proliferation. The function of BA during *in vitro* shoot multiplication is to break the apical dominance, stimulate growth of new shoots [15].

**Table 1:** Effect of sterilant on *in vitro* establishment of cv. Sai Sharbati explants.

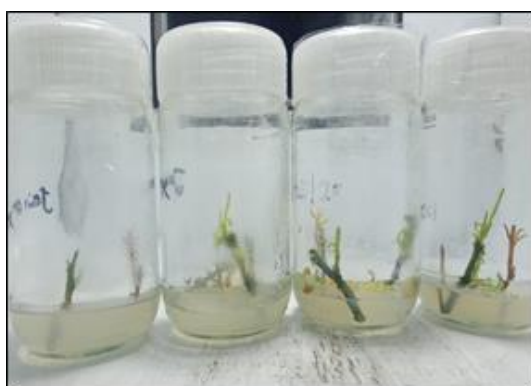
Sterilants		Time (min)	Survival (%)	Bud sprouting (%)
HgCl <sub>2</sub> (%)	NaOCl (%)			
0.05	-	3	0.0 (0.0)	0.0 (0.0)
0.05	-	5	0.0 (0.0)	0.0 (0.0)
0.1	-	3	28.00 (31.93)	49.16 (44.50)
0.1	-	5	64.43 (53.36)	74.80 (59.84)
-	10	5	0.0 (0.0)	0.0 (0.0)
-	10	10	19.00 (25.82)	4.36 (12.04)
-	15	5	26.63 (31.05)	17.70 (24.86)
-	15	10	49.33 (44.60)	30.63 (33.59)
<b>CD<sub>0.05</sub></b>			1.08 (0.71)	1.08 (0.82)
<b>SE±</b>			0.35 (0.23)	0.35 (0.27)

Values in parenthesis are arc sine transformed values. CD = Critical difference; SE = Standard error

**Table 2:** Effect of plant growth regulators on *in vitro* shoot multiplication of cv. Sai Sharbati.

Medium Composition MS (Basal medium) + GR (mg/l)			Multiplication rate	Average length (cm) shoot	Average no. of leaves
BA	Kin	GA <sub>3</sub>			
0.1	-	1.0	1: 1.5	3.13	4.20
0.5	-	1.0	1: 2.8	4.53	5.03
1.0	-	1.0	1: 2.4	3.63	4.33
1.5	-	1.0	1: 2.5	3.40	4.60
2.0	-	1.0	1: 1.8	3.50	4.46
-	0.1	1.0	1: 1.4	2.76	4.30
-	0.5	1.0	1: 1.5	3.30	4.16
-	1.0	1.0	1: 1.9	3.46	4.20
-	1.5	1.0	1: 1.3	3.63	4.13
-	2.0	1.0	1: 1.6	3.53	3.96
<b>CD<sub>0.05</sub></b>				0.59	0.46
<b>SE±</b>				0.20	0.15

Values in parenthesis are arc sine transformed values. CD = Critical difference; SE = Standard error

**Fig 1:** *In vitro* establishment of cv. Sai Sharbati explants.**Fig 2:** *In vitro* shoot multiplication of cv. Sai Sharbati.

**Table 3:** *In vitro* rooting of cv. Sai Sharbati.

Strength	Concentration of growth regulators (mg/l)		Days taken for root initiation	Percent rooting	Root length (cm)	No. of roots per shoot
	NAA	IBA				
Full strength	0.5	-	17	10.23 (18.64)	1.03	1.30
	1.0	-	20	30.23 (33.34)	1.23	1.46
	1.5	-	18	40.36 (39.40)	1.76	1.80
	2.0	-	22	20.26 (26.74)	1.53	1.63
	-	0.5	19	40.43 (39.46)	2.16	1.73
	-	1.0	19	60.23 (50.88)	2.83	2.63
	-	1.5	21	50.56 (45.30)	2.46	2.43
	-	2.0	20	40.30 (39.39)	2.26	2.13
half strength	0.5	-	18	15.50 (23.17)	2.10	1.43
	1.0	-	20	20.16 (26.67)	2.56	1.63
	1.5	-	21	40.46 (39.48)	2.36	2.13
	2.0	-	20	40.40 (39.44)	2.06	1.80
	-	0.5	16	50.40 (45.21)	3.10	2.16
	-	1.0	13	80.16 (63.52)	3.83	3.43
	-	1.5	15	60.16 (50.84)	3.00	3.00
	-	2.0	17	40.26 (39.37)	2.83	2.60
CD <sub>0.05</sub>				0.63 (0.40)	0.51	0.66
SE±				0.22 (0.14)	0.17	0.23

Values in parenthesis are arc sine transformed values. CD = Critical difference; SE = Standard error

**Table 4:** Hardening of *in vitro* raised plantlets of cv. Sai Sharbati

Potting mixture	Survival percentage	Average no. of leaves	Average plant height (cm)
Cocopeat	90.23 (71.79)	6.43	6.46
Sand	10.20 (18.61)	4.30	4.46
Cocopeat : sand (1:1)	40.30 (39.38)	4.53	5.36
Cocopeat : perlite (1:1)	50.56 (45.30)	4.50	5.16
Cocopeat : red soil (1:1)	60.46 (51.02)	4.60	5.86
Cocopeat : red soil : FYM (1:1:1)	70.00 (56.76)	5.43	6.16
CD <sub>0.05</sub>	1.15 (0.97)	0.53	0.70
SE±	0.36 (0.30)	0.16	0.22

Values in parenthesis are arc sine transformed values. CD = Critical difference; SE = Standard error



**Fig 3:** *In vitro* rooting of cv. Sai Sharbati



**Fig 4: Hardened plantlets of cv. Sai Sharbati in cocopeat**



**Fig 5: Hardened plantlets in big polybags.**

#### ***In vitro* rooting:**

Various researchers reported that auxin promotes good quality of roots. Popularly IBA and NAA used as *in vitro* root induction in different citrus species [16]. For *in vitro* rooting, the shoots were separated and transferred individually to full and half strengths of MS medium fortified with various concentration of IBA and NAA. Incorporation of 1.0 mg<sup>l</sup><sup>-1</sup> IBA resulted in 80.16 % rooting (Table-3; Fig-3). Average 3.43 roots regenerated from each shoot on half strength MS medium. Our results are in contrast to Paudyal and Haq [17] who found that NAA was superior to IBA for *in vitro* root induction in Pummelo when shoots were transferred into half strength MS medium.

#### **Hardening and acclimatization:**

Rooted plantlets obtained from previous experiments were removed from agar gelled medium and transplanted in different potting mixture (Table-4). Physical, chemical and biological properties of potting mixture are greatly influence on *in vitro* produced plantlets. It was observed that out of different potting mixture tried, cocopeat showed highest survival (90.23) with maximum plant height 6.46 cm (Fig-4). These results are in compliance with various workers who reported cocopeat to be the best potting mixture with high survival percentage in different tissue culture raised plants of various species [18-20]. This may be due to better aeration, water holding and nutrient supplying capacity of cocopeat as compared to sand [21]. The plants were kept in the glasshouse with controlled environmental condition and then transferred to polybags.

## **Conclusion**

*In vitro* propagation of cv. Sai Sharbati would ensure bulk production of true to type, disease free planting material throughout year. Out of different growth regulators tried BAP (0.5 mg<sup>l</sup><sup>-1</sup>) combination with GA<sub>3</sub> (1.0 mg<sup>l</sup><sup>-1</sup>) showed superior results with multiplication rate of 1:2.8. *In vitro* raised shoots were best rooted on half strength MS medium supplemented with IBA (1.0 mg/L). Hardening of *in vitro* plantlets was done in different potting mixtures, from which highest survival was observed on cocopeat.

#### **Acknowledgments:**

Authors are very thankful to management and technical staff, Genuine Biotech, Osmanabad, Maharashtra (India) for providing tissue culture laboratory facilities, financial assistance and continuous encouragement to carry out this study.

**Conflicts of interest:** The authors stated that no conflicts of interest.

## **References**

1. Gmitter FG and Hu X. The possible role of Yunnan, China, in the origin of contemporary Citrus species (Rutaceae). *Economic Botany*, 1990, 44 (2): 267-277.
2. Scora RW. On the history and origin of citrus. *Bulletin of the Torrey Botanical Club*, 1975, 102(6): 369-375.
3. Anonymous. 2019. Area and production of horticulture crops- All India. 2018-2019. [www.agricoop.nic.in](http://www.agricoop.nic.in)

4. Kour K and Singh B. *In vitro* multiplication of rough lemon (*Citrus jambhiri* Lush.). *IOSR Journal of Agriculture and Veterinary Science*, 2012, 1(4): 05-09.
5. Narang N and Jiraungkoorskul W. Anticancer Activity of Key Lime, *Citrus aurantifolia*. *Pharmacognosy reviews*, 2016, 10(20): 118-122.
6. Rathore JS, Rathore MS, Singh M, Singh RP and Shekhawat NS. Micropropagation of mature tree of *Citrus lemon*. *Indian journal of biotechnology*, 2007, 6(2): 239-244.
7. Shivankar VJ and Rao CN. Biological control of citrus insect pests. In: Singh S, Nagvi SAMH (Eds) *Citrus* (1<sup>st</sup> ED), International Book Distribution Co., Lucknow, India, 2001, pp. 347-367.
8. Raja WH. Studies on micropropagation in citrus rootstocks and effects of arbuscular mycorrhizal inoculation on *in vitro* raised plants. Ph.D. thesis, IARI, Division of Fruit & Horticultural Technology, New Delhi, India, 2012.
9. Murashige T and Skoog F. A revised medium for rapid growth and bio assay with tobacco tissue cultures. *Physiologia Plantarum*, 1962, 15: 473-497.
10. Murashige T. Plant propagation through tissue cultures. *Annual review of plant physiology*, 1974, 25: 135-166.
11. Gomez KA and Gomez AA. Statistical procedures for agricultural research. *John wiley and Sons*, New York. 1984, pp. 357- 427.
12. Rana JS, Singh R. *In vitro* clonal propagation of Kagzi lime (*Citrus aurantifolia* Swingle) through shoot tips. *Progressive Horticulture*, 2002, 34(1): 27-34.
13. Beura S, Singh R and Jagadev PN. *In vitro* multiplication studies in gladiolus cv. American Beauty. *Orissa Journal of Horticulture*, 2003, 31, pp. 101-105.
14. Sujata U, Syamal MM and Hamidullah I. Micropropagation of sweet orange cv. Mosambi through shoot tips and nodal segments. *Indian Journal of Horticulture*, 2010, 67(4): 21-25.
15. Muna AS, Ahmad AK, Mahmud K and Rahman KA. *In vitro* propagation of a semi-dwarfing cherry rootstock. *Plant Cell, Tissue and Organ Culture*, 1999, 59(3): 203-208.
16. Carimi F and De Pasquale F. Micropropagation of citrus. In: Jain SM and Ishii K (eds.) *Micropropagation of woody trees and fruits*. Kluwer, Netherlands, 2003, pp. 589-619.
17. Paudyal KP and Haq N. *In vitro* propagation of pummelo (*Citrus grandis* L. Osbeck). *In Vitro Cell, Developmental Biology of Plants*, 2000, 36(6): 511-516.
18. Bharati K, Prasad M, Mir H and Pal. *In vitro* regeneration and acclimatization of banana cv. Malbhog. *Current Journal of Applied Sciences and Technology*, 2018, 31(4): 1-6.
19. Chabukswar MM and Deodhar M. Rooting and hardening of *in vitro* plantlets of *Garcinia indica* Chois. *Indian journal of Biotechnology*, 2005, 4(3): 409-413.
20. Manjusha AVM and Sathyanarayana BN. Acclimatization studies in stevia (*Stevia rebaudiana* Bert.). In: 4<sup>th</sup> international symposium on acclimatization and establishment of micropropagated plants, 8<sup>th</sup>-12<sup>th</sup>, December, 2008, Bangalore, Abstracts, 37p.
21. Prabhuling G, Mastiholi AB and Kerutagi MG. Low-cost gelling agents for micropropagation of banana (*Musa acuminata*) cv. 'grande naine'. *International Journal of Plant Sciences*, 2014, 9(1): 46-51.